## The Roaming Atom: Straying from the Reaction Path in Formaldehyde Decomposition

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We present a combined experimental and theoretical investigation offormaldehyde (H<sub>2</sub>CO) dissociation to H2 and CO at energies just above the threshold forcompeting H elimination. High-resolution state-resolved imaging measurements of the CO velocity distributions reveal two dissociation pathways. The first proceeds through awell-established transition state to produce rotationally excited CO and vibrationally cold H2. The second second dissociation pathway yields rotationally cold CO in conjunctionwith highly vibrationally excited H 2. Quasi-classical trajectory calculations performed ona global potential energy surface for H 2CO suggest that this second channel represents anintramolecular hydrogen abstraction mechanism: one hydrogen atom explores large regions of the potential energy surface before bonding with the second H atom, by-passing the saddle point entirely.